PATENT COOPERATION TREATY

rom the NTERNAT	IONAL SEARCH	IING AUTH	ORITY		
NIERNATIONAL SEARCHING AUTHORITY To: DAVID R. MARSH ARNOLD & PORTER LIP STREET, N.W. WASHINGTON, D.C., DC 20004				PCT WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY	
				INTERNATIO	
					(PCT Rule 43bis.1)
				Date of mailing (day/month/year)	0 6 NOV 2007
	s or agent's file re	ference		FOR FURTHER	ACTION See paragraph 2 below
19025.021	-1 11 21 N		T- 7 - 7 - 10 - 10 - 10 - 10 - 10 - 10 -	1	
PCT/US04	al application No.		International filing date		Priority date (day/month/year)
		ation (IPC)	28 June 2004 (28.06.200 or both national classification		24 May 2004 (24,05,2004)
IPC(8): 0	C12Q 1/68(2006. 35/6,69.1,320.1,3	01);C07H 21	/04		
Applicant	133/0,09.1,320.1,3	23;350/330;	130/23.3		
PTC THE	RAPEUTICS				
1. This o	pinion contains in	dications rela	ating to the following item	s:	
\boxtimes	Box No. I	Basis of the	opinion		
\Box	Box No. II	Priority			
Ē	Box No. III	Non-establi	shment of opinion with re	gard to novelty, inve	ntive step and industrial applicability
\boxtimes	Box No. IV		ty of invention	,	эт ч төр най начовай аррачасто,
Box No. IV Lack of unity of invention Box No. V Reasoned statement under Rule 43bir, 1(a)(i) with regard to novelty, inventive step or industrial applicability, citations and evoluntations supporting such statement.			o novelty, inventive step or industrial		
	Box No. VI	Certain doc	uments cited		
	Box No. VII	Certain defe	cts in the international app	olication	
	Box No. VIII	Certain obs	ervations on the internation	nal application	
2, FUR	THER ACTION	N			
If a demand for international preliminary examination is made, this opinion will be considered to be a written opinion of the International Preliminary Examining Authority ("IPEA") except that this does not apply where the applicant chooses an Authority other than this one to be the IPEA and the chosen IPEA has notified the International Bureau under Rule 66.1bin(b) that written opinions of this International Searching Authority will not be so considered.					
If this opinion is, as provided above, considered to be a written opinion of the IPEA, the applicant is invited to submit to the IPEA a written reply together, where appropriate, with amendments, before the expiration of 3 months from the date of mailing of Form PCITISA/220 or before the expiration of 22 months from the priority date, whichever expires later. For further options, see Form PCITISA/220,					
3. For further details, see notes to Form PCT/ISA/220.					
Name and mailing address of the ISA/ US Mail Stop PCT, Ann: ISA/US Commissioner for Patents P.O. Box 1450 Facsimier Moraling, Virginia 22313-1450 Facsimier Moraling,			15 October 200	tion of this opinion 7 (15.10.2007)	Authorized officer Stephanie K. Mujnmerf, Ph.D. Telephone No. 571-272-0872
	,				

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No.	
PCT/US04/20751	

Box N	o. I Basis of this opinion
1. With	regard to the language, this opinion has been established on the basis of:
\bowtie	the international application in the language in which it was filed
	a translation of the international application into, which is the language of a translation furnished for the purposes of international search (Rules 12.3(a) and 23.1(b)).
2. With inven	regard to any nucleotide and/or amino acid Sequence disclosed in the international application and necessary to the claimed tion, this opinion has been established on the basis of:
a,	type of material
	a sequence listing
	table(s) related to the sequence listing
b.	format of material
	on paper
	in electronic form
c.	time of filing/furnishing
	Contained in the international application as filed.
	filed together with the international application in electronic form.
	furnished subsequently to this Authority for the purposes of search.
	in manual subsequentity to this Authority for the purposes of search.
3. 🖂	In addition, in the case that more than one version or copy of a sequence listing and/or table(s) relating thereto has been filled or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filled or does not go beyond the application as filled, as appropriate, were furnished.
. Additi	ional comments:

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No.
PCT/US04/20751

	101100010101
Box No. IV Lack of unity of invention	
In response to the invitation (Form PCT/ISA/206) to pay additional fees paid additional fees under protest and, where applicable, the pre paid additional fees under protest but the applicable protest fee not paid additional fees ander protest but the applicable protest fee not paid additional fees	otest fee
This Authority found that the requirement of unity of invention is not co	emplied with and chose not to invite the applicant to
pay additional fees. 3. This Authority considers that the requirement of unity of invention in accordance.	
complied with	nee with Rate 15.1, 15.2 and 15.5 is
not complied with for the following reasons:	
See the lack of unity section of the International Search Report(Form PCT/ISA	v/210)
· ·	
	•
•	
4. Consequently, this opinion has been established in respect of the following parts	of the international application:
all parts.	
the parts relating to claims Nos. 1-27	

Form PCT/ISA/237 (Box No. IV) (April 2005)

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No. PCT/US04/20751

INTERNATIONAL SEARCHING AUTHORITY				
Box No. V Reasoned statement under Rule 43 bis.1(a)(i) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement				
1. Statement				
Novelty (N)	Claims NONE	YES		
	Claims 1-27	_NO		
Inventive step (IS)	Claims NONE	_YES		
	Claims 1-27	_NO		
7.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1				
Industrial applicability (IA)	Claims 1-27	_YES		
	Claims NONE	_NO		
2. Citations and explanations:				
Please See Continuation Sheet				

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No. PCT/US04/20751

Supplemental Box	
In case the space in any of the preceding boxes is not sufficient.	

V. 2. Citations and Explanations:

Claim Interpretation

The term in an absence of SEQ ID NO.4* is being given the broadest reasonable interpretation in light of the specification. The term is not explicitly defined in the spec. Interest, SEQ ID NO.4 is released to as NREP and as "SEQ ID NO.4 sets forth a NREP, a 336 nucleotide region of a YEOF SUTE." B. 5 of specification and it is noted that searching the sequence against unacleotide databases does not necessarily provide art where the sequence is deleted. However, the nucleotide boundaries of SEQ ID NO.4 are not established relative to the context of the overall fill-length YEOF 5 UTT. Therefore, without clear nucleotide boundaries of the region comprising SEQ ID NO.4, the term is being interpreted as reading on art where the 5' UTR is deleted partially, either at the 5' end of the UTR, the 3' end of the UTR of from the middle.

The term UTR having a NeRP1 (SEO ID NO. 4) is also being given the broadest reasonable interpretation in light of the specification. As noted above, the limitations of SEQ ID NO. 4 are not clearly defined. The term is being interpreted as the opposite of in the absence of SEQ ID NO.4 and is interpreted as treading on any theory a full length VEOF 5 UTR is present in the mulciotide

The limitations of SIQ ID NO.3 are also not explicitly defined in the spot. Instead, SIQ ID NO.3 is referred to as PTCDE! and as "SEQ ID NO.3 est forth at PTCRE!; a 70° moteroids region of VERF STLTE" (b. 6 of predictation) and like SIQ ID NO.4, the nucleotide boundaries of SIQ ID NO.3 are not established relative to the full-length 5 'UTR and sarryling the sequence against nucleotide databases does not necessarily provide at where the sequence is deleted. Therefore, the term when the FTCRE is no SIQ ID NO.3' is being interpreted as reading on art the term is being interpreted as reading on at where the 5' UTR is prairially deleted, either at the 5' end of the UTR, it is 7 and the UTR is 10° SIQ in the sequence comprising SIQ ID NO.3, a fragment thereof, or a complement of either' is being interpreted are reading on any where the SIQ ID NO.3, a fragment thereof, or a complement of either' is being interpreted as reading on any where the SIQ ID NO.3, a fragment thereof, or a complement of either' is being interpreted as reading on any where the SIQ ID NO.3.

Claims I-27 tack novelty under PCT Article 33(2) as being anticipated by Forsythe et al. (Molecular and Cellular Biology, 1996, vol. 16, no. 9, p. 4604-4613). Forsythe teaches a method of analyzing the effect of hypoxia inducible factor on the expression of YEOF (Abstract).

With regard to claims I-10 and 14, Forsythe teaches a variety of nucleic acid constructs and nucleic acids that comprise a nucleic acid encoding a reporter polypeptide, wherein the nucleic acid acquence encoding a reporter polypeptide is operably linked to a NeRP, said NeRP (SEQ 10 NOs), and expression of acid reporter polypeptide is capable of being modulated relative to in an absence of said NeRP (Figure I.A, where a variety of construct) comprising deficiency on the CPU Teach (see Teach 10 Nos), and and therefore in this absence of SEQ 10 Nos); see allow, a Acid, Sec. J. Typoptic 1 passing constructs.

Form PCT/ISA/237 (Supplemental Box) (April 2005)

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No. PCI/US04/20751

Supplemental Box

In case the space in any of the preceding boxes is not sufficient.

heading, where the UTR is linked to luciferase reporter gene).

With regard to claims 11-13 and 15-21, Forsythe teaches a reporter construct wherein said VEGF 9 UTR is in an absence of SEQ ID NO34 and contains an intrino (Figure IA, where a variety of constancts comprising deletions of the 5 UTR, and therefore in the absence of SEQ ID NO34, see also p. 4605, ool. 1, "eporter plasmid construct" heading, where the UTR is inteed either to VEGF ORF or to be of the total content of the property plasmid constructs the adming, where the UTR is linked either to VEGF ORF or luciferase reporter gaze), and are produced in vitro (p. 4605, where the constructs are produced in vitro (p. 4605, where the constructs are produced in vitro (p. 4605, where the constructs are

With regard to claims 22-24, Forsythe teaches a nucleic acid molecule that comprises 95-99% sequence identity with a nucleic acid molecule of SEQ DN NO.3 a fragment thereof or a complement of either, consists of SEQ DN NO.3 or a fragment or complement thereof, or consists of a nucleic acid linked to a reporter polypeptide wherein the nucleic acid sequence consists of SEQ DN NO.3 (Figure 1A, where a variety of constructs comprising deletions and full length versions, see KpnI of the 5' UTR, and therefore comprising SEQ DN NO.3).

With regard to claims 23-27, Forsythe teaches a nucleic acid molecule that comprises 95-99% sequence identity with a nucleic acid molecule of SEQ ID NO-4, a fragment thereof or a complement of either, consists of SEQ ID NO-4 or a fragment or complement thereof, or consists of a nucleic acid infact to a reporter polypeptide wherein the nucleic acid sequence consists of SEQ ID NO-6 (Figure 1A, where a variety of constructs comprising deletions and full length versions, see RpnI of the 5' UTR, and therefore comprising SEQ ID NO-4).

Claims 22-27 lack novelty under PCT Article 33(2) as being anticipated by Kamiya et al. (US Patent 6,057,437; May 2000) teach the specific nucleotide sequences of VEGF 3' and 5' UTR regions (Table I, col., 10).

With regard to claims 22-24, Kamiya teaches a nucleic said molecule that comprises 95-99% sequence identity with a nucleic acid molecule of SSEQ DNO-3 a fragment thereof a complement of either, consists of SEQ DNO-3 are fragment for complement and their, consists of SEQ DNO-3 are fragment of complement thereof, or consists of a nucleic acid linked to a reporter polypeptide wherein the nucleic acid sequence consists of SEQ DNO-3 (Table 1, col. 10, etc. sequence alignment be blow).

٩	2y	1	TCCAGAGAGAAGTCGAGGAAGAGAGAGAGAGAGAGAGAGA	60
3	Эb	337	TCCAGAGAGAAGTCGAGGAAGAGAGAGAGAGAGAGAGAGA	396
(QΥ	61	AGCGAAAGCGACAGGGGCCAAAGTGAGTGAGCTGCTTTTTGGGGGTGACCGCCGGAGCGCGCG	120
3	Ob	397	AGCGAAAGCGACAGGGGCAAAGTGAGTGACCTGCTTTTTGGGGGTGACCGCCGGAGCGCGG	456
9	Py.	121	CGTGAGCCCTCCCCCTTGGGATCCCGCAGCCAGTCGCGCTGACGGACAGACA	180
1	Ob	457	CGTGAGCCCTCCCCCTTGGGATCCCGCAGCTGACCAGTCGCGCTGACGGACAGACA	516
٩	Эy	181	GACACOGCCCCAGCCCCAGCTACCACCTCCCCCGGCCGGCCGGGCCAGCACAGTGGACGCG	240
1	Ob	517	GACACOGCCCCAGCCCCAGCTACCACCTCCCCCGGCCGGCGGGCGG	576
(⊋у	241	GCGGCGAGCCGGGGCAGGGCCCGAGCCCGCCCGAGGCGGGGTGGAGGGGTCGAGGGTCGGG	300
3	Olo	577	GCGGCGAGCCGGGGCCGGAGCCCGGAGCCCGGAGGCGGGTGGAGGGGTCGGG	636
9	2y	301	GCTCGCGGCGTCGCACTGAAACTTITCGTCCAACTTCTGGCTGTTCTCGCTTCGGAGGA	360
ī	Ob	637	GCTCGCGGCGTCGCACTGAAACTTTTCGTCCAACTTCTGGGCTGTTCTCGCTTCGGAGGA	696
(βY	361	GCCGTGGTCCGCGCGGGGAGCCGAGCCGAGCCGAGCAGTCCTAGCTCGGGC	420
Į	Olo	697	GCCGTGGTCCGCGCGGGGGAAGCCGAGCCGAGGAGCCGCGAGAAGTGCTAGCTCGGGC	756
9	2y	421	CGGGAGGASCCGCAGCCGGAGGAGGAGGAGGAGGAGAGAGA	480
Ī	Ob	757	CGGGAGGAGCCGCAGCCGGAGGAGGAGGAGGAGAAGAAGA	816
(ЗУ	481	CCGCAGTGGCGACTCGGCGCTCGGAAGCCGGGCTCATGGACGGGTGAGGCGGCGGTGTGC	540
Į	Ob de	817	CCGCAGTGGCGACTCGGCGCTCGGAAGCCGGGCTCATGGACGGGTGAGGCGGCGGTGTGC	876
9	ЭY	541	GCAGACAGTGCTCCAGCCGCGCGCGCCCCCAGGCCCTGGCCCGGGCCTCCGGCCGG	600
I	0b	877	GCAGACAGTGCTCCAGCCGCGCGCTCCCCAGGCCCTGGCCCGGGCCTCGGGCCGGGAA	936
(ЗУ	601	GGAAGAGTAGCTCGCCGAGGCGCGGAGAGCGGGCCGCCCCCACAGCCCGAGCCGGAGA	660
I	do	937	GGAAGAGTAGCTCGCCGAGGCGCCGAGAGAGCCGGCCCCCACAGCCCGAGCCGGAGA	996

WRITTEN OPINION OF THE INTERNATIONAL SEARCHING AUTHORITY

International application No. PCT/US04/20751

Supplemental Box

In case the space in any of the preceding boxes is not sufficient.

With regard to claims 23-27, Kam'ay teaches a mudici acid molecule that comprises 95-99% sequence identity with a nucleic acid molecule of SSQ ID MO-4, a fragment thereof or complement of clience, consists of SSQ ID MO-4 or fragment or complement thereof, or consists of a nucleic acid linked to a reporter polypeptide wherein the nucleic acid sequence consists of SSQ ID MO-4 (Table 1, col. 10, see sequence alignment below).

```
1 TCGCGGAGGCTTGGGGCAGCCGGGTAGCTCGGAGGTCGTGGCGCTGGGGGCTAGCACCAG 60
            DЪ
         1 TCGCGGAGGCTTGGGGCAGCCGGGTAGCTCGGAGGTCGTGGCGCCTGGGGGCTAGCACCAG 60
Qν
        61 CGCTCTGTCGGGAGGCGCAGCGGTTAGGTGGACCGGTCAGCGGACTCACCGGCCAGGGCG 120
        61 CGCTCTGTCGGGAGGCGCAGCGGTTAGGTGGACCGGTCAGCGGACTCACCGGCCAGGGGG 120
Db
       121 CTCGGTGCTGGAATTTGATATTCATTGATCCGGGTTTTATCCCTCTTCTTTTTCTTAAA 180
Qv
       121 CTCGGTGCTGGAATTTGATATTCATTGATCCGGGTTTTATCCCCTTCTTTTTTCTTAAA 180
рь
Oν
       Db
       Οv
       241 CTTGAATCGGGCCGACGGCTTGGGGAGATTGCTCTACTTCCCCAAATCACTGTGGATTTT 300
       241 CTTGRATCGGGCCGACGGCTTGGGGAGATTGCTCCTACTTCCCCAAATCACTGTGGGATTTT 300
Db
Qy
       301 GGAAACCAGCAGAAAGAGGAAAGAGGTAGCAAGAGC 336
       301 GGAAACCAGCAGAAAGAGGAAAGAGGTAGCAAGAGC 336
Dh
```

Claims 1-27 meet the criteria set out in PCT Article 33(4), and thus have industrial applicability because the subject matter claimed can be made or used in industry.

Form PCT/ISA/237 (Supplemental Box) (April 2005)